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AMENDMENTS TO LB 756

1	1.	Strike	the	original	sections	s and	insert	the	fol	lowing
2	new sections:									
3	"Sa	ction 1	ı	Sections	1 to 8 c	of thi	ie act	ahall	he	known

- 3 "Section 1. Sections 1 to 8 of this act shall be known
- 4 and may be cited as the Cancer Drug Repository Program Act.
- 5 Sec. 2. For purposes of the Cancer Drug Repository
- 6 Program Act:
- 7 (1) Cancer drug means a prescription drug used to treat
- 8 (a) cancer or (b) the side effects of a prescription drug used to
- 9 treat cancer;
- 10 (2) Department means the Department of Health and Human
- 11 Services Regulation and Licensure;
- 12 (3) Health care facility has the definition found in
- 13 section 71-413;
- 14 (4) Health clinic has the definition found in section
- 15 71-416;
- 16 (5) Hospital has the definition found in section 71-419;
- 17 (6) Pharmacy has the definition found in section 71-425;
- 18 (7) Physician office means the office of a physician or
- 19 an osteopathic physician;
- 20 (8) Prescribing practitioner means a health care
- 21 practitioner licensed under the Uniform Licensing Law who is
- 22 authorized to prescribe cancer drugs; and
- 23 (9) Prescription drug has the definition found in section
- 24 71-1,142.

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- Sec. 3. The department shall establish a cancer drug
- 2 repository program for accepting donated cancer drugs and
- 3 dispensing such drugs to Nebraska residents. Participation in the
- 4 program shall be voluntary.
- 5 Sec. 4. Any person or entity, including but not limited
- 6 to, a cancer drug manufacturer or health care facility, may donate
- 7 cancer drugs to the cancer drug repository program. Cancer drugs
- 8 may be donated at a physician office, pharmacy, hospital, or health
- 9 clinic that elects to participate in the program and meets criteria
- 10 established by the department for such participation.
- 11 Sec. 5. (1) A cancer drug shall only be accepted or
- 12 dispensed under the cancer drug repository program if such drug is
- 13 in its original, unopened, sealed, and tamper-evident unit dose
- 14 packaging, except that a cancer drug packaged in single unit doses
- 15 may be accepted and dispensed if the outside packaging is opened
- 16 but the single-unit-dose packaging is unopened.
- 17 (2) A cancer drug shall not be accepted or dispensed
- 18 under the cancer drug repository program if (a) such drug bears an
- 19 expiration date that is earlier than six months after the date the
- 20 drug was donated or (b) such drug is adulterated or misbranded as
- 21 described in section 71-2401 or 71-2402.
- 22 (3) Subject to limitations provided in this section,
- 23 unused cancer drugs dispensed under the medical assistance program
- 24 established in section 68-1018 may be accepted and dispensed under
- 25 the cancer drug repository program.
- 26 Sec. 6. (1) A physician office, pharmacy, hospital, or
- 27 health clinic that accepts donated cancer drugs under the cancer

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- 1 drug repository program shall comply with all applicable provisions
- 2 of state and federal law relating to the storage, distribution, and
- 3 dispensing of such drugs and shall inspect all such drugs prior to
- 4 dispensing to determine if they are adulterated or misbranded as
- 5 described in section 71-2401 or 71-2402. Such drugs shall only be
- 6 dispensed pursuant to a prescription issued by a prescribing
- 7 practitioner. Such drugs may be distributed to another
- 8 participating physician office, pharmacy, hospital, or health
- 9 clinic for dispensing.
- 10 (2) A physician office, pharmacy, hospital, or health
- 11 clinic may charge a handling fee for distributing or dispensing
- 12 cancer drugs under the cancer drug repository program. Such fee
- 13 shall be established in rules and regulations adopted and
- 14 promulgated by the department. Cancer drugs donated under the
- 15 program shall not be resold.
- 16 Sec. 7. (1) Any person or entity which exercises
- 17 reasonable care in donating, accepting, distributing, or dispensing
- 18 cancer drugs under the Cancer Drug Repository Program Act or rules
- 19 and regulations adopted and promulgated under the act shall be
- 20 immune from civil or criminal liability or professional
- 21 disciplinary action of any kind for any injury, death, or loss to
- 22 person or property relating to such activities.
- 23 (2) The donation of a cancer drug by a cancer drug
- 24 manufacturer does not absolve the manufacturer of any criminal or
- 25 civil liability that would have existed but for the donation,
- 26 including, but not limited to, liability for failure to transfer or
- 27 communicate product or consumer information or the expiration date

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- 1 of the donated cancer drug.
- Sec. 8. The department, with the approval of the Board
- 3 of Pharmacy, shall adopt and promulgate rules and regulations to
- 4 carry out the Cancer Drug Repository Program Act. Initial rules
- 5 and regulations under the act shall be adopted and promulgated no
- 6 later than ninety days after the operative date of the act. Such
- 7 rules and regulations shall include, but are not limited to:
- 8 (1) Eligibility criteria and other standards and
- 9 procedures for physician offices, pharmacies, hospitals, and health
- 10 clinics that accept and distribute or dispense donated cancer
- 11 drugs;
- 12 (2) Necessary forms for administration of the cancer drug
- 13 repository program, including, but not limited to, forms for use by
- 14 persons or entities that donate, accept, distribute, or dispense
- 15 cancer drugs under the program;
- 16 (3) The maximum handling fee that may be charged by
- 17 physician offices, pharmacies, hospitals, or health clinics that
- 18 accept and distribute or dispense donated cancer drugs; and
- 19 (4)(a) Categories of cancer drugs that the cancer drug
- 20 repository program will accept for dispensing and (b) categories of
- 21 cancer drugs that the program will not accept for dispensing and
- 22 the reason that such drugs will not be accepted.
- 23 Sec. 9. This act becomes operative on September 15,
- 24 2003.".